

# Structural injury underlying mottling in ponderosa pine needles exposed to ambient ozone concentrations in the San Bernardino Mountains near Los Angeles, California

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**Abstract** For several decades, southern California experienced the worst ozone pollution ever reported. Peak ozone concentrations have, however, declined steadily since 1980. In this study, the structural injuries underlying ozone symptoms in needles of ponderosa pine (*Pinus ponderosa* Dougl. ex Laws.) collected in summer 2006 from one of the most polluted sites in the San Bernardino Mountains were investigated using serial sections examined by light and electron microscopy. Ozone-specific light-green diffuse mottling was observed in the current-year needles, whereas older foliage showed brownish mottling similar to winter fleck injury. Especially, within the outer layers of mesophyll, many markers of oxidative stress, typical for ozone, were observed in both apoplast and symplast. Altogether within cells of mottles, these markers were indicative of hypersensitive-like response, whereas degenerative structural changes were diagnosed in the surrounding mesophyll. Evidence of drought stress and frost injury to older needles was also detected. Hence, mottling injury appeared to be primarily caused by ozone stress, however, other environmental stressors also determined the symptom morphology and distribution, especially within the older foliage.

**Keywords** Ponderosa pine · Los Angeles basin · Ozone visible injury · Microscopic diagnosis · Needle histochemistry · Mesophyll ultrastructure

## Introduction

Air pollution levels and the effect on the mountain forests of southern California

For more than 50 years, southern California has experienced some of the highest air pollution ever recorded with ozone (O<sub>3</sub>) and nitrogen compounds as the airborne pollutants of greatest concern (Bytnerowicz et al. 2007, 2008; Takemoto et al. 2001). From the 1970s to the present, the June–September 24 h mean O<sub>3</sub> concentration has dropped from 100 ppb with occasional ~600 ppb peaks to 60–70 and ~180 ppb peaks due to the implementation of effective air pollution control measures. However, with AOT40 exposure indices reaching 55–75 ppm h at the most polluted places, southern California still reports the highest ozone levels in the northern hemisphere. The Mediterranean climate and rugged topography—with mountain ranges immediately downwind of the photochemical smog source areas trapping masses of polluted air—contribute significantly to the high amount of air pollution similar to other regions in the world also experiencing elevated levels of O<sub>3</sub> (Dalstein and Vas 2005; de Bauer and Hernandez-Tejeda 2007; Diaz-de-Quijano et al. 2009). Multi-decadal nitrogen (N) deposition caused mainly by elevated concentrations of nitric acid (HNO<sub>3</sub>), ammonia (NH<sub>3</sub>) and nitrogen oxides (NO<sub>x</sub>) have caused N saturation at the most exposed Californian forest sites (Bytnerowicz and Fenn 1996; Fenn et al. 2008). Although not phytotoxic, N antagonistic and synergistic effects with O<sub>3</sub> have had

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further detrimental consequences for southern California forests (Grulke et al. 1998; Grulke and Balduman 1999).

Plant damage caused by photochemical smog has been known since the 1950s in southern California and the causal role of ozone, regarding leaf injury, was established experimentally in the 1960s (Miller et al. 1963; Takemoto et al. 2001). Further research has documented: (1) the extent of injury in California forests and the slight reduction following abatement in ozone concentration, (2) alteration of needle physiology with reduction of photosynthetic exchanges and acceleration of needle senescence, (3) alteration in whole-tree biomass and carbon retention in aboveground organs which critically increases tree susceptibility to drought, windthrow and pests, (4) overall negative effects on radial growth and (5) interactions with other stressors such as N deposition, pest outbreak and stand disturbance by forest fires (Grulke et al. 2009; Karnosky et al. 2007; Miller and McBride 1999; Takemoto et al. 2001).

### Ozone injury in conifer foliage

Under ozone stress, conifer needles show visible injury in the form of homogeneous (Hartmann et al. 2007) band-like (Flagler and Chappelka 1995; Miller and Evans 1974) needle discoloration or mottling (Sanz and Calatayud 2012; Stolte 1996) which vary in intensity and morphological details depending on the ozone dose and tree species. In pine species, chlorotic mottling is the most frequent injury reported and this symptom is one of the principal parameters used to calculate the Forest Pest Management (FPM) and Ozone Injury Index (OII) in forest health surveys in California and other parts of the US (Arbaugh et al. 1998; Grulke 2003). Morphological traits of mottle characteristics of O<sub>3</sub> stress used for diagnosis versus other biotic (e.g. insect, mite or fungal injury) and abiotic stressors include: (1) diffuse edges, (2) connection to stomata and (3) distribution on the light-exposed side of needles in the sun crown (Günthardt-Goerg and Vollenweider 2007; Miller et al. 1997). Interactions with abiotic stress factors, noteworthy frost in the case of the wintering symptomatic foliage of pines in mountain forests, complicate the diagnosis. Symptoms designated as “winter fleck” have been described and their morphological characteristics do show similarities to O<sub>3</sub>-induced mottling (Hartmann et al. 2007; Miller and Evans 1974; Stolte 1996). N compounds and oxidants other than ozone do not induce mottling (Davis 1977; Padgett et al. 2009; Takemoto et al. 2001).

Structural changes to be found in living as well as dead cells form markers of cellular physiological processes and responses (Fink 1999; Günthardt-Goerg and Vollenweider 2007; Jones 2000; Levine et al. 1996) and can be used for the diagnosis of ozone damage (Kivimäenpää et al. 2005;

Vollenweider et al. 2003a, b). Several studies have analyzed structural injuries triggered by O<sub>3</sub> stress in conifer needles with (Table 1) or without (Anttonen and Karenlampi 1996; Evans and Leonard 1991; Kainulainen et al. 2000) visible symptoms. In trees showing mottling but generally independent of visible injury, they have documented characteristic changes to the mesophyll structure, whereas other tissues remained nearly asymptomatic. Concerning the microscopic changes underlying mottling, however, little has been published so far. Indeed, these symptoms are difficult to process in view of microscopical analysis because of their (1) poor visibility, further degrading during sample fixation and processing and (2) minute size. Hence, unless special care is taken during sample handling and sectioning, the structural injuries causing mottling are easily missed. Structural injury reported so far for Jeffrey (*Pinus jeffreyi* Grev. & Balf.) or ponderosa pine (*Pinus ponderosa* Dougl. ex Laws.) dates back to the early studies using needle material showing massive visible injury as a consequence of the high doses of ozone which occurred at that time, but which no longer presently exist in California (Table 1).

### Study objectives

The aim of this study was to characterize the microscopic changes and the processes leading to mottling injury in pine needles in response to ambient O<sub>3</sub> pollution, as currently occurring in southern California. Needles from ponderosa pine, a pine species with well-documented photosynthetic sensitivity to O<sub>3</sub> (Coyne and Bingham 1982) and showing mottling both in the field (Stolte 1996) as well as in controlled conditions (Richards et al. 1968), were sampled near Camp Paivika (CP), a long-term air pollution/forest health monitoring station in the San Bernardino Mountains near Los Angeles, California, USA (Miller and Rechel 1999). Structural injury underlying mottling was investigated using transmitted light, fluorescence and electron microscopy (EM) and related to specific O<sub>3</sub> or other interacting stress effects (Günthardt-Goerg and Vollenweider 2007; Vollenweider et al. 2003a).

## Materials and methods

### Study site, climate and air pollution

The CP study site was selected as being one of those consistently displaying the severest ozone injury in the San Bernardino Mountains (Miller and McBride 1999) with OII in 1997, amounting to 57.1 versus 14.6 at an eastern SBM site (authors' unpublished data). It is located directly to the east (ca. 70 km) of Los Angeles at 1,580 m a.s.l. on the

**Table 1** Structural injury triggered by O<sub>3</sub> stress in conifer needles showing mottling or other types of necrotic flecks

Species	O <sub>3</sub> treatment/O <sub>3</sub> exposure	Needle sampling	Injury		Histological	Cytological		Reference
			Macroscopic	Microscopic				
<i>P. ponderosa</i> , <i>jeffreyi</i> , <i>coulteri</i> , <i>lambertiana</i>	350 ppb, 9-h day <sup>-1</sup> , 55 days <sup>a</sup>	Targeted at visible injury	n.i. (chlorotic mottling)		Initial injury in outer layers of mesophyll, with or without association with substomatal areas, progressing inwardly to finally encompass all mesophyll layers	Gradual loss of cell integrity		Evans and Miller (1972a)
<i>P. ponderosa</i>	450 ppb, 12-h day <sup>-1</sup> , 35 and 8 days <sup>a</sup>	Targeted at visible injury	Chlorotic mottling		Initial injury in outer layers of mesophyll, independent of substomatal areas, progressing inwardly and leading to tissue collapse concomitant to cell death.	Cell death and collapse within 5 days		Evans and Miller (1972b)
<i>Picea abies</i>	n.i.	Targeted at visible injury	Minute necrotic spots		Necrosis and transition area in mesophyll	Necrosis: disrupted cell content, condensation of the cell remnants along cell walls, coarse granular particles in cytoplasm and chloroplasts  Transition area: degenerated cytoplasm, coarse granular particles in cytoplasm, chloroplasts with a smaller size and modified shape showing membrane ruptures and stroma degeneration, degenerating mitochondria		Sutinen (1986)
<i>P. abies</i>	50–98 ppb, 7-h day <sup>-1</sup> , 113–183 days during 3 years <sup>a</sup>	Independent of visible injury	Chlorotic mottling		Initial injury in outer layers of mesophyll facing the sky progressing inwardly and downwardly to mesophyll layers facing the ground	Independent of mottling: decrease in the chloroplast and starch grain size, increased stroma condensation, reduced amount of thylakoid membranes, increased amounts of plastoglobuli, disruption of peroxysomes and mitochondria		Sutinen et al. (1990)
<i>Pinus ellioti</i>	55–71 ppb, 7-h day <sup>-1</sup> , 17–83 days <sup>a</sup>  ≤20 ppb <sup>b</sup>	Targeted at visible injury	Chlorotic mottling		Injury in mesophyll	Cell plasmolysis and cytorhysis		Evans and Fitzgerald (1993)
<i>Pinus rigida</i>	50–200 ppb, 8-h day <sup>-1</sup> , 13 weeks <sup>a</sup>	Independent of visible injury	Chlorotic needle tips, curvature at mid-needle		Injury starting in outer layers of mesophyll, around the substomatal cavities and progressing towards inner mesophyll and vascular bundle, accumulation of intercellular debris in mesophyll, degeneration of endodermis and phloem collapse	Condensation of the chloroplast stroma, thylakoid swelling, increased plastoglobuli density		McQuattie and Schier (1993)

Table 1 continued

Species	O <sub>3</sub> treatment/O <sub>3</sub> exposure	Needle sampling	Injury	Histological	Cytological	Reference
			Macroscopic			
<i>Pinus halepensis</i>	150–600 ppb, 7–12-h day <sup>-1</sup> , 2–16 days <sup>a</sup>	Independent of visible injury	Light-yellowish flecks on >50 % of the needle	Injury primarily in outer layers of mesophyll and around the substomatal cavities	Protoplasm disruption, decrease in the chloroplast and starch grain size, increased stroma condensation, thylakoid swelling and membrane disruption, increased fatty acid saturation	Anttonen et al. (1995)
<i>Pinus sylvestris</i>	100–600 ppb/12–8.4 ppm h, 2–28 days <sup>a</sup>	Independent of visible injury	With higher O <sub>3</sub> exposure only, development of large necrotic flecks in >50 % of foliage	Most severe injury in outer layers of mesophyll and around the substomatal cavities	Mesophyll cell collapse at high O <sub>3</sub> concentration, decrease in the chloroplast and starch grain size, increased stroma condensation and granulation, increase in the plastoglobuli density at low and decrease at high O <sub>3</sub> concentration	Holopainen et al. (1996)
<i>Pinus taeda</i>	47–98 ppb, 12-h day <sup>-1</sup> , 140 days <sup>a</sup>	Independent of visible injury	Chlorotic mottling, water-soaked bands, necrotic spots	n.i.	Alteration of the plasmalemma and chloroplast envelope, decrease in the chloroplast and starch grain size, increased stroma condensation, thylakoid swelling, increase in the density of plastoglobuli	Anttonen et al. (1996)
<i>Abies religiosa</i>	>110 ppb, 4–5-h day <sup>-1</sup> , 300 days <sup>b</sup>	Independent of visible injury	Whitish stippling apart from the mid-rib coalescing into reddish flecks and developing abaxially prior to needle yellowing and shedding	Discrete necrotic groups of cells with the most severe injury in outer layers of mesophyll	Disrupted cell content, condensation of the cell remnants along cell walls, plasmolysis	Alvarez et al. (1998a)
<i>P. sylvestris</i>	14.5–33 ppm h (AOT40), 20 weeks <sup>a</sup>	Independent of visible injury	Chlorotic mottling and yellowing of the previous needle generation	n.i.	Increased frequency of cytoplasmic ribosomes, transient increase in the condensation of stroma	Utriainen and Holopainen (1998)
<i>P. halepensis</i>	20 ppm h (AOT40) <sup>b</sup>	Including visible injury	Chlorotic mottling	Injury especially in outer layers of mesophyll, around the substomatal cavities and in the vascular bundle	Decrease in the chloroplast size, increased stroma condensation, increased plastoglobuli density, plastoglobuli excretion towards the vacuole, increase in amounts of tannin, starch and lipid	Soda et al. (2000)
<i>Pinus cembra</i>	n.i.	Including visible injury	Photobleaching and mottling	Discrete necrotic groups of cells in mesophyll	Disrupted cell content, condensation of the cell remnants along cell walls, accumulation of proanthocyanidins	Vollenweider et al. (2003a)

Table 1 continued

Species	O <sub>3</sub> treatment/O <sub>3</sub> exposure	Needle sampling	Injury		Histological	Cytological	Reference
			Macroscopic	Microscopic			
<i>P. halepensis</i>	35.4–56.5 ppb/ 15.5–36.7 ppm h (AOT40) <sup>b</sup>	Independent of visible injury	Chlorotic mottling		Injury in mesophyll	In association with mottling: change in the vacuolar tannin structure, decrease in the chloroplast size and frequency, increased plastoglobuli density, increased amounts of peroxysomes and mitochondria, reduced amounts of starch and lipids	Kivimäenpää et al. (2010)

n.i. no indication

<sup>a</sup> Material exposed to O<sub>3</sub>-enriched air during fumigation experiments

<sup>b</sup> Material exposed to ambient O<sub>3</sub> levels

crestline within a mixed conifer stand with a western aspect. The climate at CP is temperate to cold, sub-humid and as such, representative of a Mediterranean and mountain area (Walter and Lieth 1967; Fig. 1a). Eighty percent of the annual precipitation is deposited in October–March (Takemoto et al. 2001), mainly in the form of snow and there is a summer drought between mid-April and October. In 2006, with an annual mean temperature of 13 °C and only 717 mm precipitation, CP experienced drier climatic conditions in comparison to averages over the last 50 years, and a more severe summer drought with a precipitation deficit lasting from July to the end of the year (Fig. 1b vs. a).

Bedrock at CP is granitic and soils are coarse-loamy, mixed, mesic, and typical Haploxerolls (Arkley 1981). Forest stands at CP form the western end of the mixed conifer zone in the San Bernardino Mountains. Vegetation type is mixed conifer forest with ponderosa pine and California black oak (*Quercus kelloggii* Newb.) as the most abundant tree species at the site. In 2006, only an open stand remained over a portion of the study plot as a consequence of increased tree mortality following the drought 2002–2004 and a fire in October 2003, which burned a part of the CP site.

Ozone concentrations for the southern California region were obtained from the California Air Resources monitoring network equipped with Thermo Environmental Model 49 UV absorption monitors. Data were presented as the highest 8 h average concentration during the entire year 2006. O<sub>3</sub> concentrations for CP were obtained from the Crestline air monitoring site located 4 km to the east. O<sub>3</sub> exposure indices were calculated using the Ozone Calculator program (William Jackson, USDA Forest Service, <http://webcam.srs.fs.fed.us>). The SUM00 index is an exposure dose resulting from multiplying all hourly concentrations by time (h). The SUM06 index was derived by multiplying the number of hours when O<sub>3</sub> concentrations were >60 ppb (0.06 ppm) by those concentrations. These two indices were calculated for 24 h for a 4-month period (June 1–September 30). In comparison with the European studies, AOT40 exposure index was also calculated as a cumulative dose of values >40 ppb (0.04 ppm) for the standardized daylight hours (08:00–20:00 PST) during the 6-month growing season (April 1–September 30).

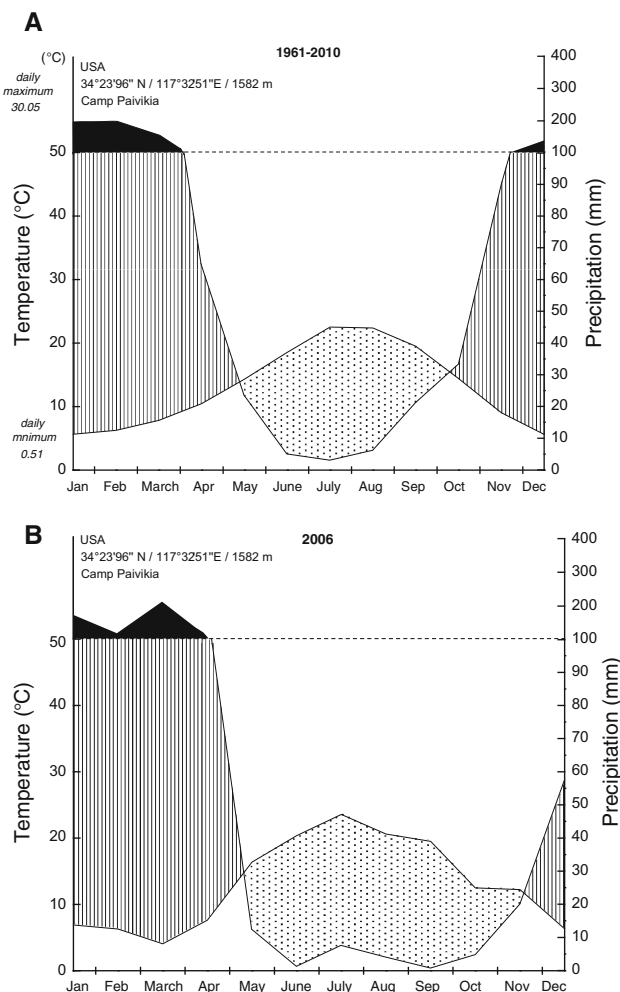
#### Sampling and microscopic analysis

Sampling occurred on the afternoon of September 10, 2006 and focused on mottling symptoms typical of O<sub>3</sub> stress (Sanz and Calatayud 2012), whilst deliberately excluding material showing other types of injury (Vollenweider and Günthardt-Goerg 2006). One tree displaying representative mottling injury on accessible (1–2.5-m high) branches



exposed to full sunlight was selected. Ten current (C) and ten 1-year-old (C + 1) symptomatic needles were collected along the same branch segments. Older needle generations were not harvested because of the risk of confusion with other stress factors. Symptomatic middle needle segments were trimmed and immediately fixed using 100 % methanol and regular or EM-grade 2.5 % glutaraldehyde—the latter two solutions buffered at pH 7.0 with a 0.067 M Soerensen phosphate buffer—and stored at 4 °C for 12 days. At the Swiss Federal Institute for Forest, Snow and Landscape Research (WSL) laboratory, glutaraldehyde fixed samples were entirely infiltrated with renewed fixing solution by evacuation before storage at

4 °C until further processing. To investigate histological, cytological and histochemical changes in mottling symptoms and surrounding needle parts, samples were processed in three different ways: (1) methanol-fixed samples were hand-microtomed to obtain thick cuttings (30 µm); (2) samples fixed with regular glutaraldehyde were dehydrated with 2-methoxyethanol (three changes), ethanol, n-propanol, and n-butanol (Feder and O'Brien 1968), embedded in Technovit 7100 (Kulzer HistoTechnik) and semi-thin cuttings (2 µm) were sectioned using a Reichert 2050 microtome; (3) EM-grade fixed samples were post-fixed in buffered 2 % OsO<sub>4</sub>, dehydrated by a series of graded ethanol, infiltrated by a series of graded propylene oxide/Epon 812 mixture (with DDSA, NMA and DMP hardener), embedded in Epon and ultra-thin cuttings (70 nm) were sectioned using a Reichert UltraCut S ultra microtome. For each type of preparation, batches of serial cross-sections were cut through symptomatic needle segments (about 650 µm of needle tissue per batch) and cuttings at symptom core locations were sorted out. Thick and semi-thin sections were stained with different methods (Table 2), mounted in water or DPX and observed using a Leica microscope Leitz DM/RB, with 5× to 100× objectives and diascopic light illumination. Thin sections were mounted on copper grids and contrasted using saturated uranyl acetate in 50 % ethanol and lead citrate (Reynolds procedure). Sections were observed using a Philips CM 12 transmission electron microscope.



**Fig. 1** Climate diagram summarizing the climatic conditions at Camp Paivikia during the 1961–2010 reference period (a) and in 2006 (b). Diagrams are plotted according to Walter and Lieth (1967). Between 0 and 100 mm precipitation monthly, 20 mm on the right ordinate is equal 10 °C of average temperature on the left ordinate. Above 100 mm (dashed line), precipitation is plotted using a scale 5 times larger. The dry and moist seasons are outlined with a dotted line and dashed/solid area, respectively. Mean yearly temperature and precipitation: 13.04 °C and 926 mm between 1961 and 2010 (a), 13.03 °C and 717 mm in 2006 (b)

## Results

### Air pollution

In 2006, CP was among the most O<sub>3</sub> polluted sites in southern California with the highest 8 h average concentration of 0.143 ppm (Fig. 2). There were clearly pronounced seasonal differences in O<sub>3</sub> concentrations with low values of the pollutant in the October–March winter season and much higher levels during the summer photochemical smog season (April–September, Fig. 3). Accordingly, average O<sub>3</sub> concentrations in June, July and August were the highest (0.062, 0.059 and 0.061 ppm, respectively). The maximum hourly value of 0.164 ppm at CP was recorded in July, while the mid-winter maximum hourly concentration dropped to 0.050 ppm. The annual average O<sub>3</sub> concentration was 0.042 ppm. The O<sub>3</sub> exposure indices were: SUM00–161.9 ppm h, SUM06–103.3 ppm h, and AOT40–58.34 ppm h. These results confirm the high concentrations of O<sub>3</sub> and its exposure indices at CP. Although the recorded concentrations were lower than in 2002–2005 (figure not shown), they were still higher than other comparable forest sites in North America and Europe (Bytnerowicz et al. 2008).

### Visible symptoms in C and C + 1 needles

In C needles from CP, O<sub>3</sub>-like visible injury was observed in the form of tiny rounded light green mottles scattered on the unshaded abaxial as well as adaxial needle side exposed to full sunlight. The mottle's average diameter  $\pm$  SE was  $221 \pm 20 \mu\text{m}$  (range 121–305  $\mu\text{m}$ ,  $n = 10$  symptoms measured using five needles, Fig. 4a, b). These symptoms developed around the stomata amid needle tissue showing a lighter green color than shaded needle parts and they had diffuse boundaries. They were observed on most ponderosa pines at the study site, but were limited to the C needle generation. Contrastingly, in adjacent C + 1 and older needle generations, larger light brownish mottles encompassing several stomatal lines and with sharp boundaries were observed. Their average diameter  $\pm$  SE was  $543 \pm 67 \mu\text{m}$  (range 317–926  $\mu\text{m}$ ,  $n = 10$  symptoms measured using five needles, Fig. 4c). They were observed on the abaxial and adaxial needle side and their frequency increased with the needle age and full sunlight illumination. Compared to C needles, the C + 1 needles showed a darker green background color.

### Needle structure of C and C + 1 needles

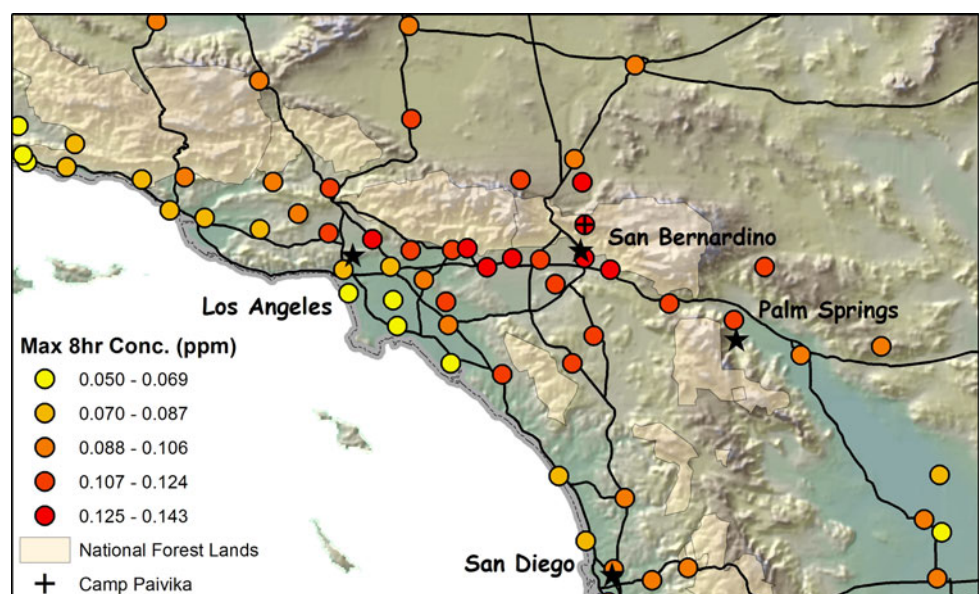
In the mesophyll outside of mottles, little vitality differences between cells were observed, but structural gradients were evident (i.e. from the inner to outer mesophyll: an increase in the chloroplast condensation and proanthocyanidin oxidation and an apparent reduction in the chloroplast and starch grain size; figures not shown). Assimilative cells throughout mesophyll were functional as indicated by (1) thin and mostly unlignified cell walls (Figs. 5c, 7d, k),

(2) uncondensed cytoplasm with low amounts of lipid droplets (Figs. 5e, 6a, 7g), (3) large and uncondensed nuclei (Figs. 5c, g, 7d, i), (4) chloroplasts with distinct grana, few plastoglobuli and little starch despite sampling in the afternoon (Fig. 6i), (5) peroxysomes and mitochondria without apparent injury and occasional smooth endoplasmic reticulum (Fig. 6a), (6) vacuoles with undisrupted tonoplast and without large lipid accumulation or autophagic remains (figure not shown). However, both inside and outside of mottles, an enlarged periplasm was occasionally observed. In asymptomatic cells, the chloroplasts were small, their stroma generally condensed and the membranes poorly resolved (Fig. 6a, i). Tannins in the form of proanthocyanidins were observed in cell walls, nuclei and vacuoles (Figure not shown). In C + 1 versus C needles, the vacuolar proanthocyanidin deposits had their frequency strongly increased and their shape changed from mostly droplet-like to an homogeneous or finely granular and only occasionally ribbon-like or spongy form (Figs. 7a, d vs. 5a, c, 6a). Other tissues showed a healthy structure and biotic injury was missing.

### Structural changes in mottles of C needles

Inside mottles of C needles, the strongest structural changes were found within the outer mesophyll cells adjacent to the substomatal chamber (Figs. 4d, 5b vs. a). Mottles included dead and degenerating cells and the transition to asymptomatic inner or surrounding mesophyll occurred within a single cell layer. At the mottle center, massive cell wall thickening obstructed most of the intercellular space (Figs. 5b, d, h, j, k, 6f). Full embedding in Technovit (Fig. 5)—respectively Epon (Fig. 6)—resin of these cell

**Fig. 2** Maximum 8-h O<sub>3</sub> concentrations throughout southern California in 2006



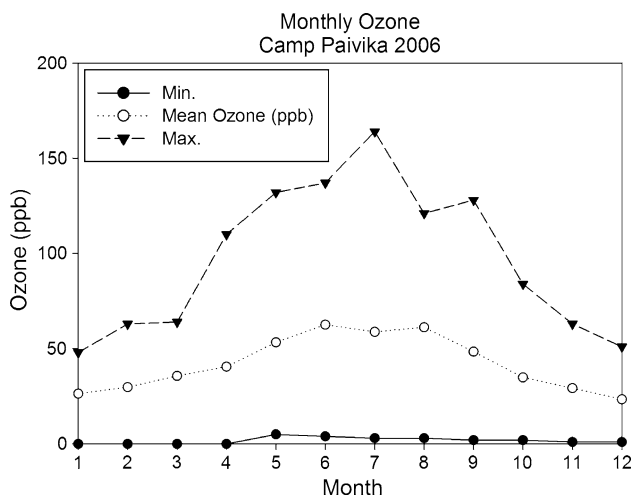
**Table 2** Staining methods (stain dilution in w/v)

Stain	Reference	Solution	Staining time (min)	Color in transmitted light	Excitation (nm)	Histochemical target	Figure
Vanilin acid <sup>a</sup>	Sarkar and Howarth (1976)	Vanilin 10 % in ethanol:HCl 37 % = 2:1	Observed in reagent	Red	–	Proanthocyanidins	Figure not shown
Toluidine blue O <sup>b</sup>	Feder and O'Brien (1968)	0.05 % aq. acetate buffer pH 4.4	10	Blue	–	Metachromatic stain	4d, e, 5a–d, 7a–f, m
Fuchsin acid <sup>b</sup>	Feder and O'Brien (1968)	1 % Fuchsin acid aq.	20	Pink	–	Metachromatic stain	4e, 7a–f, m
Coomassie brilliant blue <sup>b</sup>	Wetzel et al. (1989), modified	0.025 % in 100 % ethanol:acetic acid = 3:1	35	Blue	–	Proteins	5g, h, 7i, j
Calcofluor white Mr2 <sup>b</sup>	Munck (1989)	0.1 % in 50 % ethanol aq.	2	–	340–380	Cellulose	Figure not shown
Coriophosphine <sup>b</sup>	Weis et al. (1988)	0.03 % aq.	3	–	450–490	Pectins	7n
PARS <sup>b</sup>	Gahan (1984)	0.5 % periodic acid; Schiff reagent; 0.5 % potassium metabisulfite in 0.05 N HCl	10; 20; 3 × 5	Pink	–	Polysaccharides	5k
Aniline blue <sup>b</sup>	Gerlach (1984)	0.005 % in Soerensen buffer pH 8.2	10	–	340–380	Callose	Figure not shown
Phloro-glucinol <sup>b</sup>	Schneider (1979) as cited by Clark (1981), modified	Saturated in 18 % HCl	Observed in reagent	Pink	–	Lignin	5i, j, 7k, l
Sudan-black B <sup>b</sup>	Brundrett et al. (1991)	0.2 % in 100 % ethanol:glycerin = 1:1	Observed in reagent	Blue-black	–	Lipids	5e, f, 7g, h

Material

<sup>a</sup> Hand-microtome cuttings<sup>b</sup> Semi-thin Technovit-embedded cuttings





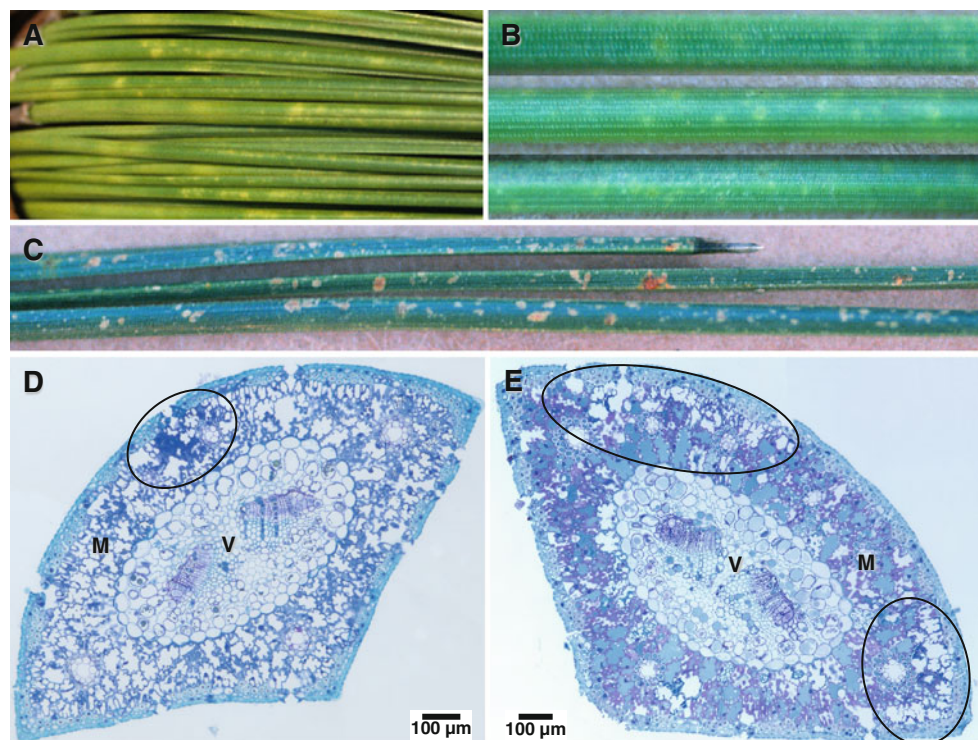
**Fig. 3** Seasonal changes (monthly mean, minimum and maximum values) in  $O_3$  concentrations at the Crestline air monitoring station in 2006

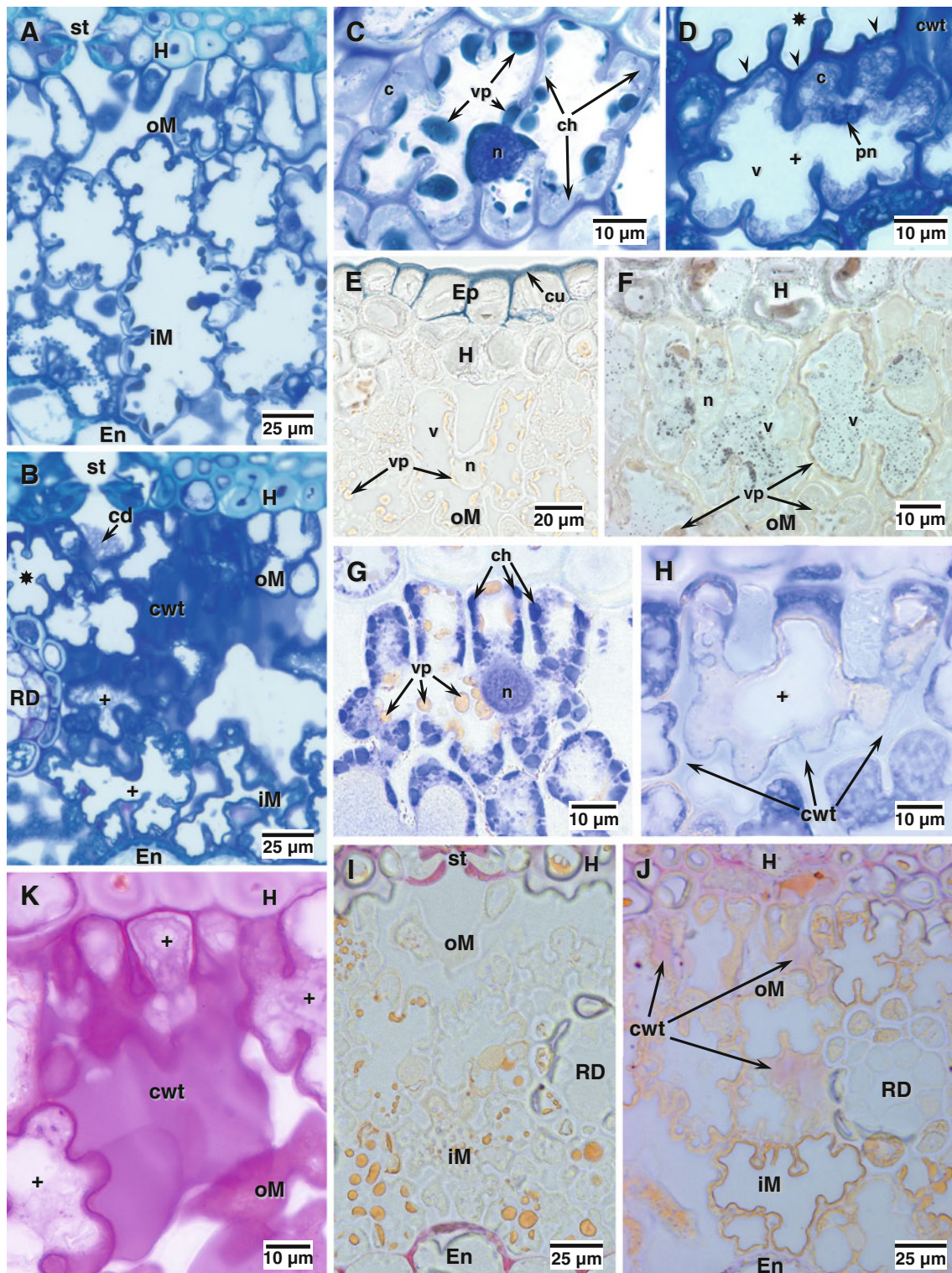
wall thickenings and adjacent cells excluded the possibility that these structures resulted from sample processing artifacts. Further, these intercellular masses showed structural traits typical for cell walls, including (1) an ultrastructure similar to middle lamella material (Fig. 6f), (2) a structural continuity with the cell wall outer layers (Figs. 5k, 6f) and (3) a typically polysaccharidic (Fig. 5k) matrix made of mostly pectins. Additionally, small protein (Fig. 5h), lignin (Fig. 5j) and cellulose (figure not shown) fractions were detected, while callose was missing. Heterogeneous and

loosely arranged cell debris, structurally and histochemically contrasting with cell wall thickening and presumably leaked from dead cells, were also observed in the intercellular space (Figs. 5b, 6f). Within dead cells, the cell lumen was empty and the condensed cell content remnants were densely packed onto cell walls (Fig. 5d). Surrounding cells showed varying degenerative features indicative of a gradual cell decompartmentation and typically evidenced by blistering of the tonoplast and other membranes (Figs. 5d, 6c). In comparison to the asymptomatic mesophyll, the cell content showed more lipid droplets in the vacuole (Fig. 5f vs. e), but a smaller protein fraction in the cytoplasm (Fig. 5h vs. g). Less vacuolar proanthocyanidins were observed than in the mesophyll outside mottles.

At a subcellular level, the cytoplasm and organelles of cells inside mottles showed advanced degeneration contrasting with the moderate changes found in the surrounding mesophyll (Fig. 6b vs. a). Injury progress in the cytoplasm was evidenced by increasing condensation and formation of coarse granular particles (Fig. 6c–e vs. 6a), lipid accumulation (Fig. 6d), empty vesicle formation (Fig. 6c, d) and tonoplast blistering (Figs. 5d, 6c). In terminal stages, only amorphous and coagulated particles with few still recognizable structures such as large starch grains remained (Fig. 6e). Regarding organelles, nuclei were small, deformed and strongly condensed (picnotic nuclei, Figs. 5d, 6g), mitochondria showed progressive disruption of the matrix and inner membranes and apparently evolved to empty and rather inflated vesicles (Fig. 6c, d, g, h), and

**Fig. 4** Macro-(a–c) and micro-morphological (d, e) characteristics of the mottling symptoms observed at Camp Paivikia in 2006. **a, b** Light-green diffuse mottling in C needles; symptoms were found on the light exposed side of needles and were centered on stomata. **c** Brownish mottling in C + 1 needles; symptoms were larger than the diffuse symptoms and showed an irregular contour. **d, e** Needle serial cross-sections through mottling symptoms. Structural injury (circle) was found in outer mesophyll (M) cells whereas veins (V) showed no injury. In C needles (d), injury was restricted to a small group of cells adjacent to stomata. In C + 1 needles (e), an extended portion of outer mesophyll was injured





peroxysomes had disappeared. Chloroplasts also showed a characteristic structural evolution including size reduction, a decrease in grana frequency and thickness, an increase in plastoglobuli density and an enhanced stroma condensation (Fig. 6j–l). During the terminal degeneration stages

(Fig. 6l), the plastoglobuli were no longer identifiable and the chloroplasts finally disrupted except for the starch grains (Fig. 6e). Degenerative processes in chloroplasts and probably also in other organelles, particularly membrane injury, could contribute to lipid droplet formation



◀ **Fig. 5** Structural (a–d) and histochemical (e–j) changes at the tissue (a–b, i–j) and cellular (c–h, k) level found inside (b, d, f, h, j, k) versus outside (a, c, e, g, i) mottling symptoms in mesophyll of C needles showing *greenish* diffuse mottling. **b** (*Mottling*) versus **a** (*no mottling*): Cells within mottled areas were dead (\*) or showed advanced degeneration (+). At mottle centre, massive cell wall thickening (*cwt*) filled the whole intercellular space. Other type of intercellular material included amorphous and loosely arranged cell debris (*cd*). **d** (*Mottling*) versus **c** (*no mottling*): Dead cells showed condensed cell content remnants appressed to cell walls (*arrow-heads*), whereas less degenerated cells showed a partly disrupted protoplasm with irregular vacuolar outline and a few still recognizable organelles (*arrows*). Nuclei had shrunk to highly condensed and deformed picnotic structures (*pn*). **f** (*Mottling*) versus **e** (*no mottling*): Lipid droplets (*dark blue*) accumulation in vacuoles of outer mesophyll cells at mottle centers. **h** (*Mottling*) versus **g** (*no mottling*): The protein cell content was decreased, whereas a small protein fraction (light blue staining) was inlaid within the massive cell wall thickenings. **j** (*Mottling*) versus **i** (*no mottling*): Lignin inlay (pink) within the massive cell wall thickenings—notice the constitutively lignified cell walls in hypodermis and endodermis cells. **k** (*Mottling*): The massive cell wall thickenings (*pink*) were primarily made of polysaccharides (pectins). Other structures: *c* cytoplasm, *ch* chloroplasts, *cu* cuticula, *En* endodermis, *Ep* epidermis, *H* hypodermis, *iM/oM* inner/outer mesophyll, *RD* resin duct, *st* stomata, *v* vacuole, *vp* vacuolar proanthocyanidins

(Fig. 6l) prior to accumulation in the cytoplasm (Fig. 6d) and coalescence in the vacuole (Fig. 5f).

#### Structural changes in mottles of C + 1 needles

Similar to C needles, the most prominent changes in mottles of C + 1 needles were observed in the outer layers of mesophyll, but they extended wider and were not restricted to a single stomatal line (Fig. 4e). Symptoms reached variable depth inside the assimilative tissue, whereas injury to other tissues was generally nonexistent (Figs. 4e, 7b, c vs. a). Cells in rows just below the hypodermis were necrotic and often broken but rarely collapsed and their cell remnants were scattered in the intercellular space (Fig. 7c). In the still remaining complete cells, cell death was indicated by an empty cell lumen presumably as a consequence of tonoplast, plasmalemma and organelle disruption and condensation of oxidized cell content remnants along cell walls (Fig. 7f). A transition zone of cells showing less advanced degenerative features surrounded the necrotic mottle center (Fig. 7b, e). A high level of nuclear material condensation was observed, indicative of chromatin breaking similar to that found after programmed cell death (Fig. 7e; Levine et al. 1996; Alvarez et al. 1998b). Histochemically, the amount of proteins in cells inside mottles was lower than in the asymptomatic mesophyll (Fig. 7j vs. i), whereas the cytoplasmic lipid content was slightly increased (Fig. 7h vs. g). The vacuolar proanthocyanidins inside mottles showed a lower frequency and a ribbon-like structure compared to homogenous filling in the surrounding mesophyll (Fig. 7b, e, vs. a, d). In cell

walls, the middle lamella showed a higher degree of lignification than was found in the surrounding mesophyll (Fig. 7l vs. k). In dead cells and in those within the transition to still healthy-looking mesophyll, thickened cell walls, structurally similar to, but not as massive as those observed in C needles, were detected (Fig. 7m). They were only found in the outer and better illuminated cells and mainly consisted of oxidized pectins (Fig. 7n), without any other histochemically detectable fraction.

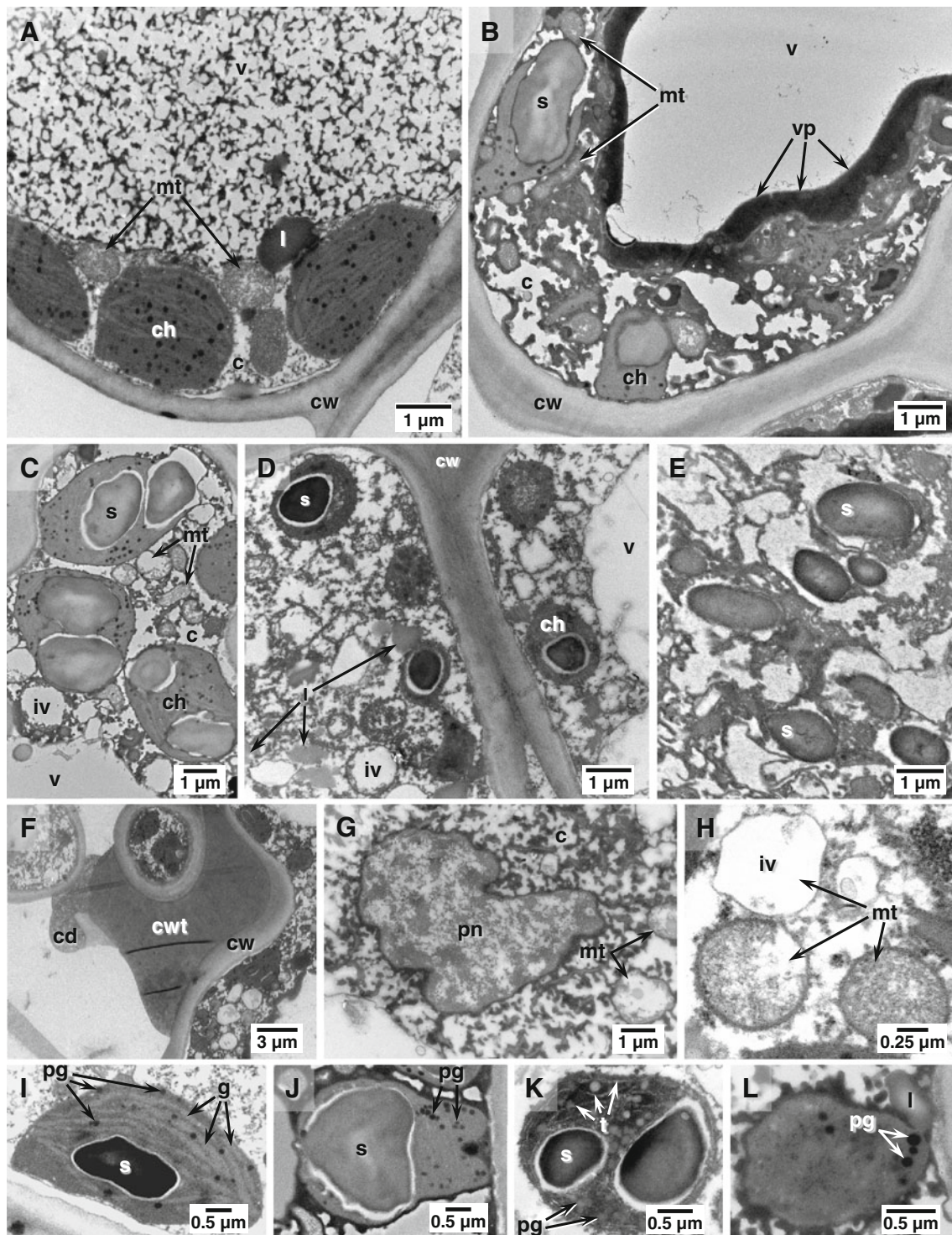
## Discussion

### Visible injury

With 58 ppm h, the CP O<sub>3</sub> exposure was much larger than the 24 ppm h threshold for O<sub>3</sub> injury in C and C + 1 needles as found during fumigation experiments of seedlings (Takemoto et al. 2001). Two types of visible O<sub>3</sub> injury differing according to the needle age were observed in the ponderosa pine needles sampled at CP. In C needles, the light green mottling was typical of O<sub>3</sub> injury with respect to its morphology, distribution and by comparison to O<sub>3</sub> injury documented in *Pinus halepensis*, *strobus* and *uncinata* (Díaz-de-Quijano et al. 2011; Kivimäenpää et al. 2010; Sanz and Calatayud 2012). Mottles found in adjacent C + 1 and older needles showed morphological traits similar to the so-called “winter fleck” injury (brownish dots with sharp edges; Hartmann et al. 2007; Miller and Evans 1974; Stolte 1996), but which were also reminiscent of chlorotic mottling by O<sub>3</sub> stress (symptom frequency increasing with needle age and higher illumination on both abaxial and adaxial needle sides) as found in *Pinus ponderosa* and other conifers (Alvarez et al. 1998a; Flagler and Chappelka 1995; Grulke 2003). With both symptoms adjacent to each other on the same branches, our results suggest that mottling in C and C + 1 needles was related ontologically. In the field; however, mottling is generally observed in C + 1 and older needles (Grulke and Lee 1997; Miller et al. 1997), whereas mottling already occurred in C needles in the examined material. This finding may relate to the higher O<sub>3</sub> exposure in the San Bernardino Mountains than in other O<sub>3</sub>-polluted regions of the world (Bytnerowicz et al. 2008).

### Microscopic symptoms in C needles

With regard to their distribution and structure, changes typical of O<sub>3</sub> stress were found within mottles and surrounding mesophyll. At the tissue level, the injury gradient, with highest severity in outer mesophyll cell layers next to stomata, together with the unscathed epidermis and vascular bundle were typical of O<sub>3</sub> stress in ponderosa pine



(Evans and Miller 1972a) and other conifers (Alvarez et al. 1998a; Anttonen and Karenlampi 1996; Günthardt-Goerg and Vollenweider 2007; Soda et al. 2000; Sutinen et al. 1990). Symptom proximity to stomata contrasted with the distal location observed in conifer needles (Alvarez et al. 1998a) and leaves from broad-leaved trees (Vollenweider

et al. 2003a) with a dorso-ventral polarity. Stomata in foliage of the latter species are primarily located on the shaded abaxial side versus stomata located all around pine needles. Whatever the species,  $O_3$  injury primarily develops within the more light-exposed portions of the mesophyll where it exceeds the cell detoxifying capacities in



◀ **Fig. 6** Ultrastructural changes in cells inside (**b–g, j–l**) versus outside (**a, h, i**) mottling symptoms in mesophyll of C needles showing greenish diffuse mottling. **b** (*mottling*) versus **a** (*no mottling*): The cell structure became increasingly disorganized in cells inside mottles. Notice in **a** the granular structure of proanthocyanidins in the vacuole (*v*). **c–e** With increasing injury, cytoplasm (*c*) condensed in coagulated particles, lipid droplets (*l*) accumulated, the chloroplast (*ch*) and mitochondria (*mt*) structure disrupted and the tonoplast boundary became irregular. In the most degenerated cells, only the starch grains (*s*) could still be recognized. **f** Cell debris (*cd*) and massive cell wall thickening (*cwt*) within the outer mesophyll apoplast. **g** Deformed and highly condensed picnotic nucleus (*pn*). The surrounding cytoplasm particles are condensed and the mitochondria showed advanced degeneration. **h** Degeneration sequence of mitochondria. The matrix and inner membrane system showed injury increasing until only a nearly empty and rather inflated vesicle (*iv*) made of the mitochondria outer membrane was left. **i–j** Degeneration sequence of the chloroplast structure. **i** In cells outside mottles, the stroma appeared condensed and the grana (*g*) poorly resolved. **j–l** Within mottles, the chloroplasts showed a size reduction and an increased plastoglobuli (*pg*) density, further stroma condensation, grana stack and thylakoid (*t*) frequency reduction and poor membrane resolution. In a few cases, the extrusion of lipids (*l*) outside chloroplasts was observed (*L*). Other structures: *cw* cell wall, *vp* vacuolar proanthocyanidins

synergy with photo-oxidative stress (Foyer et al. 1994; Günthardt-Goerg and Vollenweider 2007). Hence, the peculiar injury distribution, with regard to the stomata observed here and in other pines, should relate primarily to the concomitant location of stomata and better illuminated mesophyll cells and secondarily to the proximity to Reactive Oxygen Species (ROS) from O<sub>3</sub>-induced oxidative bursts in the apoplast (Baier et al. 2005; Matyssek and Sandermann 2003).

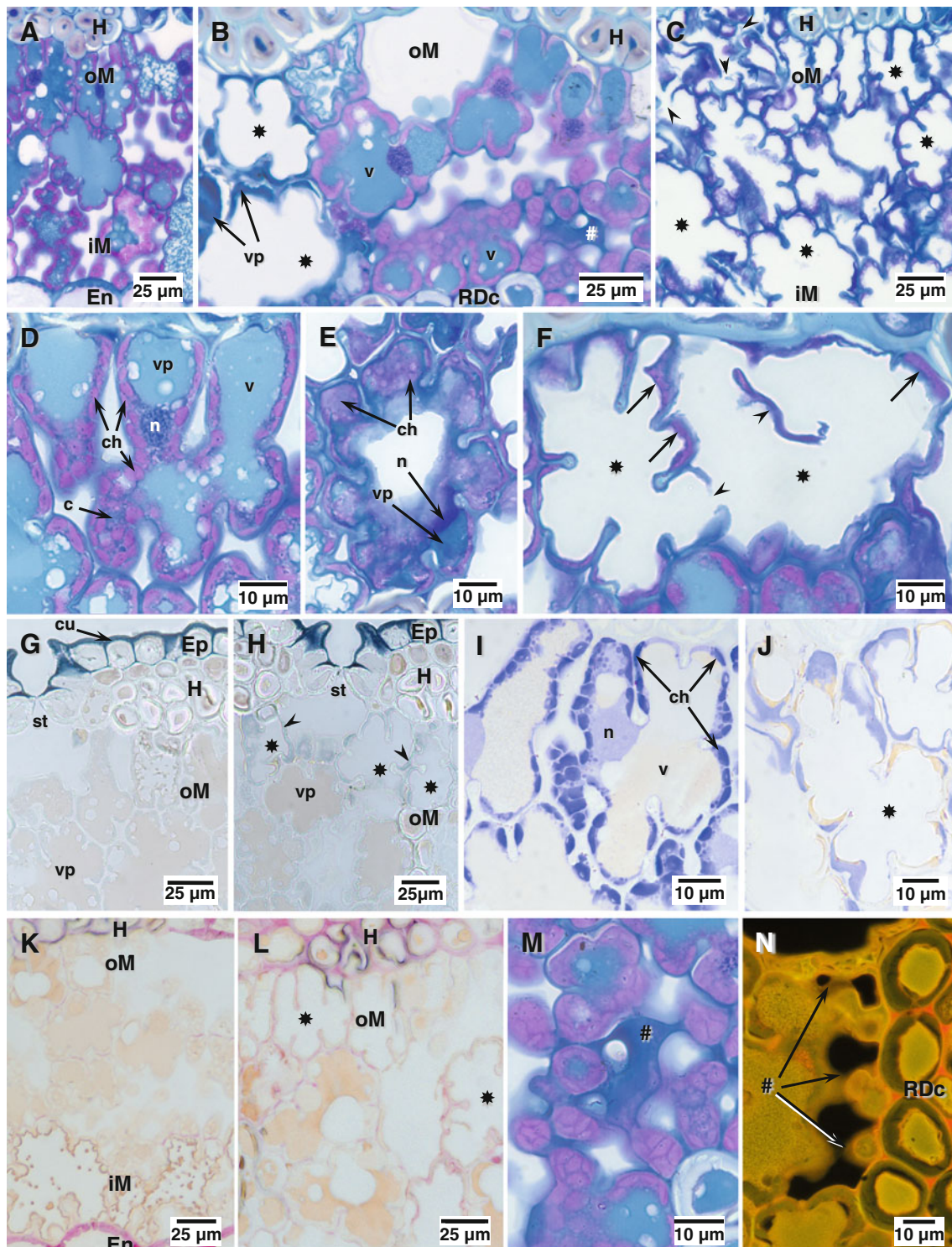
At the cellular level, changes in chloroplasts, especially those in outer cell layers, were characteristic of the effects of O<sub>3</sub> stress (Fink 1999; Holopainen et al. 1996; Kivimäenpää et al. 2005; Soda et al. 2000; Sutinen et al. 1990). Several of these changes, probably contributing to lighter green color in the unshaded needle segments, were also observed in cells outside mottles similar to the findings in O<sub>3</sub> exposed, but asymptomatic foliage (Anttonen and Karenlampi 1996; Kainulainen et al. 2000). Nevertheless, the severity of injury was higher inside than outside mottles and the local disruption process of chloroplasts appears to be the principal reason for the scattered bleached-dot morphology of visible injury. The plastoglobuli accumulation and apparent extrusion into the cytoplasm or vacuoles were indicative of the active involvement of these chloroplastic structures in lipid trafficking and thylakoid membrane turnover, as occurring during senescence and stress events (Bréhélin et al. 2007; Mikkelsen and Heide-Jorgensen 1996; Soda et al. 2000; Tevini and Steinmüller 1985), and of the aforementioned disruption process. Hence, chloroplasts are very sensitive to oxidative stress (Coyne and Bingham 1982; Foyer et al. 1994; Günthardt-Goerg and Vollenweider 2007) as a consequence of the

major ROS production occurring in this organelle in comparison to other symplastic sites (Kangasjarvi et al. 2005; Sandermann 1996; Yamasaki et al. 1997). Changes in chloroplast size and plastoglobuli density as a function of chloroplast depth gives further confirmation that O<sub>3</sub> and photo-oxidative stress promote chloroplast injury synergistically.

Among the prominent structural changes, those found in the cell walls of the outer mesophyll cells in mottle centers have never been reported for conifers, but recent observations in *Pinus uncinata* confirm such cell wall modifications as being a response to ozone stress (Díaz-de-Quijano et al. 2011). These massive thickenings were structurally and histochemically similar to characteristic markers of oxidative stress in angiosperms as a consequence of elevated O<sub>3</sub> concentrations (Bâesso Moura et al. 2011; Bussotti et al. 2005; Günthardt-Goerg et al. 1997; Vollenweider et al. 2003a) and other factors causing oxidative stress (Hermle et al. 2007). The detected protein and lignin fraction may be involved in ROS detoxification within the apoplast (Iriti and Faoro 2008; Polle 1997). Within cells from tissue showing mottling symptoms, many structural indications of terminal cell content degeneration and disruption were observed. The missing peroxysomes and degeneration sequence of the mitochondrial membrane and matrix system were particularly indicative of severe oxidative stress inside cells (Jones 2000; Richter 1993; Richter and Schweizer 1997). Besides the amorphous structures, the degradation products included lipid droplets in the cytoplasm and vacuole similar to the findings in Aleppo pines under O<sub>3</sub> stress (Soda et al. 2000) or some broad-leaved species (Mikkelsen and Heide-Jorgensen 1996). All these structural features, together with changes such as the formation of picnotic nuclei or cell content disruption inside restricted groups of mottle cells, meet the structural criteria to identify a hypersensitive-like response reminiscent of that occurring in sensitive broad-leaved trees (Günthardt-Goerg and Vollenweider 2007; Vollenweider et al. 2003a). However, these criteria have been established studying angiosperm species and confirmation from the cell physiology specifically for gymnosperms is still wanting. The occurrence of an accelerated cell senescence response (ACS) in the mesophyll surrounding mottles is suggested by markers such as oxidative injury in chloroplasts and by the accelerated needle shedding (Miller et al. 1997; Takemoto et al. 2001), although O<sub>3</sub>-induced and ontological needle aging, presumably because of oxidative stress, differ structurally (Kivimäenpää et al. 2005).

Structural traits in cells outside mottles such as periplasm enlargement, low amounts of starch and perhaps some chloroplast stroma condensation suggest a drought stress effect (Fink 1999) reflecting the environmental





conditions prior to sampling. Inside mottles, the large starch grains together with the low amounts of proanthocyanidins may indicate that the detected hypersensitive-like response occurred early in the vegetation season when still enough soil moisture was available to sustain normal gas exchanges and elevated  $O_3$  uptake (Panek 2004; Panek

and Goldstein 2001). Indeed in reference to Seigler (1998), time is needed to accumulate the costly and energy-demanding proanthocyanidins, the deposits of which are generally increased within  $O_3$ -symptomatic foliage (Kivimäenpää et al. 2010; Soda et al. 2000; Vollenweider et al. 2003a). At CP, ponderosa pine needles elongate from

◀ **Fig. 7** Structural (a–f) and histochemical (g–n) changes at the tissue (a–c, k–l) and cellular (d–j, m–n) level found inside (c, f, h, j, l, m, n) versus outside (a, d, g, i, k) mottling symptoms and transition zone (b, e) in mesophyll of C + 1 needles showing brownish mottling. c, f (Mottling) versus a, d (no mottling): Cells within chlorotic mottles were dead (\*) and cell injury increased from the inner (iM) to outer mesophyll layer (oM). Cell walls were often broken (arrowheads), especially within the outer mesophyll layer, whereas the cell lumen was empty and cell content remnants appressed to cell walls (arrows). b, e (transition zone) versus a, d (no mottling): Cells in transition zones showed local thickening of their cell walls (#) and a more or less disrupted cell content. h (Mottling) versus g (no mottling): Lipid content increase (arrowheads). j (Mottling) versus i (no mottling): Protein content decrease. l (Mottling) versus k (no mottling): Increased lignification of the cell wall by lignin (pink) primarily inlaid in the middle lamella. m, n Structure and histochemistry of cell wall thickening. Deposits were amorphous and consisted mostly of pectin as shown by their reactivity with coriphosphine. In comparison to the pectin-specific orange color, as found in the cell walls of nearby resin duct sheath cells (RDc), the brownish color of thickenings (#→) indicated an increased oxidation. Other structures: c cytoplasm, ch chloroplasts, En endodermis, Ep epidermis, H hypodermis, n nucleus, st stomata, v vacuole, vp vacuolar proanthocyanidins

April/May until the end of July during dry years (Grulke et al. 1998); In 2006, this was the period during which the highest O<sub>3</sub> concentrations were measured.

#### Microscopic symptoms in C + 1 needles

Structural injury underlying chlorotic mottling in C + 1 needles also showed typical indications of O<sub>3</sub> stress with regard to its distribution and markers of oxidative stress in apoplast and symplast. Hence, it showed the same injury pattern as found in C needles which also suggests a hypersensitive-like plant response. The necrotic cell structure and distribution was similar to that reported for C + 1 and older needles of *Abies religiosa* (Alvarez et al. 1998a) or *P. halepensis* (Kivimäenpää et al. 2010). In comparison to C needles however, several structural traits were observed only in older foliage: (1) large injuries encompassing several stomata lines, (2) broken cells, (3) less developed cell wall thickenings and (4) more disrupted and oxidized cell structures. Hence, disruptive processes in symptomatic tissues were more advanced in C + 1 than C needles which might relate to the larger time lapse since mottling establishment, as suggested by the larger amounts of proanthocyanidins in surrounding and still living versus dead mottle cells. However, the rupture and removal of entire cells were clearly not an O<sub>3</sub> effect and suggests mechanical injury by another abiotic stress factor, presumably frost (Fink 1999). The outer mesophyll location of this injury is also indicative of the aforementioned stress factor. Other indications of winter fleck injury such as symptoms in inner mesophyll and vascular bundles (Fink 1999; Miller and Evans 1974) were generally missing. In synthesis, these observations suggest an injury initially

caused by O<sub>3</sub> then aggravated during winter, presumably following the formation of ice crystals at the most damaged locations (Fink 1999). The injury was likely further intensified during the next growing season as a consequence of steady oxidative stress at the periphery of the symptomatic area. Hence, interaction between O<sub>3</sub> and frost stress is suggested by these structural changes—a process which, with the contribution of supplementary biotic and abiotic stress factors, could cause a frost hardiness reduction in the affected foliage (Fink 1999).

#### Conclusion

In conclusion, structural injury found in the apoplast and symplast of mesophyll cells underlying the mottling symptoms in ponderosa pine needles contrasted with those in surrounding areas and were indicative of severe oxidative stress suggesting hypersensitive-like responses to O<sub>3</sub>, as described in angiosperms. Other environmental stressors, i.e. drought in C and frost in C + 1 needles, contributed sizably to the morphological singularities of the symptomatic areas. Hence, structural injury underlying mottling was not only useful to gain insights about important effects of ozone on cell physiology, but also to better ascertain the diagnosis in a context of high air pollution, such as in southern California, and to distinguish the effects of other concomitant stress factors.

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